

(19) World Intellectual Property
Organization
International Bureau



(43) International Publication Date
23 June 2005 (23.06.2005)

PCT

(10) International Publication Number
WO 2005/056116 A2

(51) International Patent Classification⁷: A62D 1/00

(74) Agents: HAMER, Katherine, A. et al.; Traskbirt, Suite 300, 230 South 500 East, P.O. Box 2500, Salt Lake City, UT 84110-2550 (US).

(21) International Application Number:
PCT/US2004/040248

(22) International Filing Date: 2 December 2004 (02.12.2004)

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data:
10/727,088 2 December 2003 (02.12.2003) US

(81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SI, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.

(71) Applicant (for all designated States except US):
ALLIANT TECHSYSTEMS INC. [US/US]; M.S.
MN01-2070, 5050 Lincoln Drive, Edina, MN 55436-1097 (US).

(84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

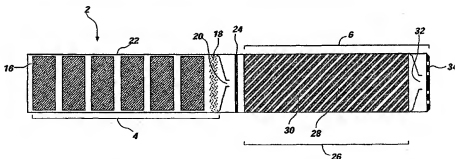
(72) Inventors: BLAU, Reed, J.; 75 East Main, Richmond, UT 84333 (US). ROZANSKI, James, D.; 589 North 600 East, Brigham City, UT 84302 (US). TRUITT, Richard, M.; 3313 North 700 East, North Ogden, UT 84414 (US). LUND, Gary, K.; 6276 South Old Highway 191, Malad, ID 83252 (US). DOLL, Daniel, W.; 740 North 2800 West, Marriott Slaterville, UT 84404-9515 (US). BRADLEY, Steven, J.; 479 East 3525 North, North Ogden, UT 84414 (US). GUYMON, Ross, W.; 122 Hillside Drive, Smithfield, UT 84335 (US). HOLLAND, John; 2321 East Combe Road, Ogden, UT 84403-5040 (US).

Published:

— without international search report and to be republished upon receipt of that report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: MAN-RATED FIRE SUPPRESSION SYSTEM



(57) Abstract: A fire suppression system for producing an inert gas mixture having a minimal amount of carbon monoxide, particulates, or smoke. The inert gas mixture may be generated by combusting a gas generant. The gas generant may be a composition that includes hexa (ammine)-cobalt(III)-nitrate. The fire suppression system also includes a heat management system to reduce a temperature of the inert gas mixture. A method of extinguishing fires is also disclosed.

-1-

MAN-RATED FIRE SUPPRESSION SYSTEM

TECHNICAL FIELD

The present invention relates to a fire suppression system. More specifically,
5 the present invention relates to a fire suppression system suitable for use in occupied or clean environments.

BACKGROUND

A fire involves a chemical reaction between oxygen and a fuel that is raised to
10 its ignition temperature by heat. The fire is extinguished by removing oxygen, reducing a temperature of the fire, separating the oxygen and the fuel, or interrupting chemical reactions of the combustion. Halogen-containing agents, such as Halon® agents, are chemical agents that have been effectively used to suppress or extinguish fires. These halogen-containing agents generate chemically reactive halogen radicals
15 that interfere with combustion processes in the fire. However, many Halon® agents, such as Halon® 1211, Halon® 1301, and Halon® 2402, have been suggested to contribute to the destruction of stratospheric ozone in the atmosphere, which has led many countries to ban their use. Therefore, effective fire fighting replacements for Halon® agents are being developed. For instance, fire suppression systems have been
20 recently developed to extinguish fires in enclosed spaces. These fire suppression systems introduce a flow of inert gas into the enclosed space to extinguish the fire. Some fire suppression systems use a source of compressed gas as the inert gas. However, the compressed gas requires a large storage area, which adds additional bulk and hardware to the fire suppression system.

25 Other fire suppression systems have utilized a propellant to generate the inert gas. The propellant is ignited to generate the inert gas, which is then used to extinguish the fire. The inert gas typically includes nitrogen, carbon dioxide (CO₂), or water. Some propellants used in fire suppression systems produce up to 20% by volume of CO₂. While CO₂ is a nonflammable gas that effectively extinguishes fires, propellants
30 that generate copious amounts of CO₂ cannot be used to extinguish fires in a human-occupied space because CO₂ is physiologically harmful. CO₂ has an Immediately Harmful to Life or Health (IDLH) value of a concentration of 4% by

-2-

volume and causes the human breathing rate to quadruple at levels from 4% by volume to 5% by volume, loss of consciousness within minutes at levels from 5% by volume to 10% by volume, and death by asphyxiation with prolonged exposure at these or higher levels. In addition, it is difficult to produce CO₂ by combustion without
5 producing significant amounts of carbon monoxide (CO), which has an IDLH of 0.12% by volume (i.e., 1200 parts per million (ppm)). Many propellants also produce other gaseous combustion products, such as ammonia (NH₃), which has an IDLH of 300 ppm; nitric oxide (NO), which has an IDLH of 100 ppm; or nitrogen dioxide (NO₂), which has an IDLH of 20 ppm. NO and NO₂ are collectively referred
10 to herein as nitrogen oxides ("NO_x"). CO₂, CO, NH₃, and NO_x are toxic to people and, therefore, producing these gases is undesirable, especially if the fire suppression system is to be used in a human-occupied space. Furthermore, many of these propellants produce particulate matter when they are combusted. The particulate matter may damage sensitive equipment, is potentially an inhalation hazard, irritates the skin and
15 eyes, and forms a hazardous solid waste that must be properly disposed of. In United States Patent No. 6,024,889 to Holland *et al.*, a chemically active fire suppression composition is disclosed. The fire suppression composition includes an oxidizer, a fuel, and a chemical fire suppressant and produces CO₂, nitrogen, and water when combusted. The composition also undesirably produces smoke and particulate matter
20 upon combustion.

Propellants based on sodium azide (NaN₃) have also been developed for use in fire suppression systems. While NaN₃-based propellants produce nitrogen as a combustion product, the propellants are problematic to produce on a large scale because NaN₃ is toxic. In addition, combusting the NaN₃ propellant produces corrosive
25 and toxic combustion products, in the form of smoke, that are very difficult to collect or neutralize before the nitrogen is used to extinguish the fire.

A nonazide-based fire suppression system is disclosed in United States Patent No. 5,957,210 to Cohrt *et al.* In the fire suppression system, ammonia is reacted with atmospheric air or compressed air to produce nitrogen and water vapor. The ammonia
30 and air are reacted in a combustion chamber of a gas turbine to produce combustion gases that are exhausted into a mixing chamber before being introduced into an enclosed space. Water is sprayed into the combustion chamber to cool the combustion

-3-

gases. The introduction of the combustion gases into the enclosed space reduces its oxygen content and extinguishes the fire.

Other fire suppression systems utilize a combination of compressed gases and propellants. In United States Patent No. 6,016,874 to Bennett, a fire extinguishing system is disclosed that uses compressed inert gas tanks and solid propellant gas generants that produce inert gases. The solid propellant gas generants are either azide- or nonazide-based and produce nitrogen or CO₂ as combustion products while argon or CO₂ are used as the compressed gases. The inert gases from each of these sources are combined to produce an inert gas having 52% nitrogen, 40% argon, and 8% CO₂ that is used to extinguish the fire.

In United States Patent No. 5,449,041 to Galbraith, an apparatus for extinguishing fires is disclosed. The apparatus includes a gas generant and a vaporizable liquid. When ignited, the gas generant produces CO₂, nitrogen, or water vapor at an elevated temperature. The hot gases interact with the vaporizable liquid to convert the liquid to a gas, which is used to extinguish the fire.

DISCLOSURE OF THE INVENTION

The present invention relates to a fire suppression system that comprises a gas generant and a heat management system. The gas generant may be formed into a pellet that is housed in a combustion chamber of the fire suppression system. Upon combustion, the gas generant pyrotechnically produces an inert gas mixture that may be used to extinguish a fire. The gas generant may produce at least one gaseous combustion product and at least one solid combustion product when combusted. The gas generant may be formulated to produce minimal amounts of toxic gases, particulates, or smoke when combusted. The inert gas mixture may comprise nitrogen and water and be dispersed from the fire suppression system within from approximately 20 seconds to approximately 60 seconds after ignition of the gas generant. The fire suppression system may also include an igniter composition that is present in powdered, granulated, or pelletized form. The igniter composition may be formed into a pellet with the gas generant.

The fire suppression system also comprises an ignition train, a combustion chamber, and an effluent train that includes the heat management system. The heat

management system cools the temperature of the inert gas mixture before the inert gas mixture exits the fire suppression system. The inert gas mixture may be cooled by flowing the inert gas mixture over a heat sink or a phase change material.

When ignited, the igniter composition may produce gaseous combustion products and solid combustion products that provide sufficient heat to ignite the gas generant. The igniter composition may be a composition including from approximately 15% to approximately 30% boron and from approximately 70% to approximately 85% potassium nitrate (known in the art as "B/KNO₃"), a composition including strontium nitrate, magnesium, and a binder ("Mg/Sr(NO₃)₂/binder"), or mixtures thereof. The gas generant may be a composition that includes hexa(amine)cobalt(III)-nitrate ("HACN"), cupric oxide (CuO), titanium dioxide (TiO₂) and polyacrylamide ([CH₂CH(CONH₂)_n) or a composition that includes HACN, cuprous oxide (Cu₂O), and TiO₂. At least one of an inorganic binder, an organic binder, or a high-surface area conductive material may also be used in the gas generant.

The present invention also relates to a method of extinguishing a fire in a space. The method comprises igniting a gas generant to produce an inert gas mixture comprising a minimal amount of carbon monoxide, carbon dioxide, ammonia, or nitrogen oxides. The inert gas mixture is then introduced into the space to extinguish the fire. The gas generant may include a nonazide gas generant composition that produces gaseous combustion products and solid combustion products. Substantially all of the gaseous combustion products produced by the gas generant may form the inert gas mixture, which includes nitrogen and water. The gaseous combustion products may be produced within from approximately 20 seconds to approximately 60 seconds after ignition of the gas generant. The solid combustion products may form a solid mass, reducing particulates and smoke formed by combustion of the gas generant. The fire may be extinguished by reducing an oxygen content in the space to approximately 13% by volume.

The gas generant may be a composition that includes HACN, CuO, TiO₂, and polyacrylamide or a composition that includes HACN, Cu₂O, and TiO₂. At least one of an inorganic binder, an organic binder, or a high-surface area conductive material may also be used in the gas generant. An igniter composition may be used to combust

-5-

the gas generant, such as a B/KNO_3 composition, a composition of $Mg/Sr(NO_3)_2$ /binder, or mixtures thereof.

BRIEF DESCRIPTION OF THE DRAWINGS

5 While the specification concludes with claims particularly pointing out and distinctly claiming that which is regarded as the present invention, the advantages of this invention can be more readily ascertained from the following description of the invention when read in conjunction with the accompanying drawings in which:

FIGs. 1 and 2 are schematic illustrations of an embodiment of a fire
10 suppression system of the present invention;

FIGs. 3a and 3b are schematic illustrations of a gas generant pellet, optionally including an igniter, usable in the fire suppression system of the present invention;

FIG. 4 is a schematic illustration of an embodiment of the fire suppression system of the present invention;

15 FIG. 5 shows the calculated mole percent of oxygen in a 100 cubic foot (2.83 cubic meter) room; and

FIGs. 6 and 7 show pressure and temperature traces of Test A and Test B.

BEST MODE(S) FOR CARRYING OUT THE INVENTION

20 A fire suppression system including a gas generating device is disclosed. The gas generating device produces an inert gas mixture that is introduced into a space having a fire. As used herein, the term "space" refers to a confined space or protected enclosure. The space may be a room or a vehicle that is occupied by humans, animals, or other living beings, or by electronic equipment. For instance, the space may be a
25 room in a residential building, a commercial building, a military installation, or other building. The space may also be a vehicle or other mode of transportation, such as an automobile, an aircraft, a space shuttle, a ship, a motor boat, a train or subway, or a race car. Since the fire suppression system may be used in a space occupied by people, the fire suppression system is "man-rated." The fire suppression system may also be
30 used in a clean environment, such as a room or vehicle that is used to store or house electronic equipment.

The inert gas mixture may be generated pyrotechnically by igniting a gas generant that produces gaseous combustion products. The gaseous combustion products may include gases that do not contribute to ozone depletion or global warming. As such, these gases may be used in the inert gas mixture. The gaseous combustion products may include minimal, nonhazardous amounts of noxious gases, such as NH_3 , CO , NO_x , or mixtures thereof. In one embodiment, the gas generant produces significantly less than the respective IDLH of each of these gases and less than 1% of an original weight of the gas generant in particulates or smoke. The gas generant may also produce minimal amounts of other carbon-containing gases, such as CO_2 . In one embodiment, the gas generant produces less than approximately 4% by volume of CO_2 . The gas generant may be formulated to produce minimal carbon dioxide, particulates, or smoke when combusted and to produce a physiologically acceptable balance of toxic gases produced under fuel rich (CO and NH_3) or fuel lean (NO_x) conditions. Solid combustion products are ultimately produced upon combustion of the gas generant and may be essentially free of products that vaporize at the flame temperature of the gas generant and may solidify upon cooling to produce particulates and smoke that are respirable.

The inert gas mixture is generated in a short time frame, so that the fire may be extinguished quickly. For instance, the gas generant may be ignited, produce the inert gas mixture, and the inert gas mixture dispersed into the space within a time frame ranging from approximately 20 seconds to approximately 60 seconds. The inert gas mixture may decrease the oxygen content in the space so that oxygen-promoted combustion reactions in the fire may be suppressed or extinguished. The inert gas mixture may also decrease the oxygen content by creating an overpressure in the space, which causes oxygen-containing gases that were present in the space to exit by a positive pressure venting system and be replaced by the inert gas mixture. The positive pressure venting system for a given space may be designed to prevent a significant overpressure in the room.

The fire suppression system 2 may include a combustion chamber 4 and an effluent train 6, as shown in FIGs. 1 and 2. The fire suppression system 2 may be formed from a material and construction design having sufficient strength to withstand pressures generated by the gas generant 8. The pressures generated in the fire

-7-

suppression system 2 may range from approximately 100 pounds per square inch ("psi") (approximately 0.690 Mega Pascals ("MPa")) to approximately 1,000 psi (approximately 6.90 MPa), such as from approximately 600 psi (approximately 4.14 MPa) to approximately 800 psi (approximately 5.52 MPa). To withstand these

5 pressures, an outer surface of the combustion chamber 4 and the effluent train 6 may be formed from a metal, such as steel. The ignition train may be electrically activated, as known in the art. The gas generant 8 and an igniter composition 14 may be housed in the combustion chamber 4. The gas generant 8 may be present in the combustion chamber 4 as a pellet 16 or the gas generant 8 and the igniter composition 14 may be

10 pelletized, as described in more detail below. Embodiments of the pellet 16 are illustrated in FIGs. 3a and 3b and are described in more detail below.

The gas generant 8 in the combustion chamber 4 may be ignited to produce the gaseous combustion products of the inert gas mixture by an ignition train using sensors that are configured to detect the presence of the fire in the space. The sensors may

15 initiate an electrical impulse in the ignition train. The sensors are conventional and, as such, are not discussed in detail herein. The electrical impulse may then ignite an initiating device 12, such as a squib, semiconductor bridge, or other conventional initiating device. Heat flux from the initiating device 12 may be used to ignite the igniter composition 14, which, in turn, ignites the gas generant 8. The igniter

20 composition 14 and the gas generant 8 are described in more detail below. When ignited or combusted, the igniter composition 14 may produce an amount of heat sufficient to ignite the gas generant 8. Alternatively, the initiating device 12 may be used to directly ignite the gas generant 8. In one embodiment, the igniter composition 14 produces solid combustion products, with minimal production of

25 gaseous combustion products. The combustion products produced by this igniter composition 14 may include a minimal amount of carbon-containing combustion products.

In addition to housing the ignition train, the combustion chamber 4 may house the igniter composition 14 and the gas generant 8. The gas generant 8 may be formed

30 into a pellet 16 for use in the fire suppression system 2. Alternatively, the pellet 16 may include the gas generant 8 and the igniter composition 14, with the igniter composition 14 present predominantly on an outer surface of the pellet 16. The gas

-8-

generant 8 may be a nonazide gas generant composition that produces gaseous combustion products and solid combustion products. The gaseous combustion products may be substantially free of carbon-containing gases or NO_x . Effluents produced by the combustion of the gas generant 8 may be substantially free of NO_2 and may have less than 100 parts per million ("ppm") of other effluents, such as CO or NH_3 . For instance, the gas generant 8 may produce nitrogen and water as its gaseous combustion products. At least a portion of the gaseous combustion products produced by combustion of the gas generant 8 may form the inert gas mixture. In one embodiment, substantially all of the gaseous combustion products form the inert gas mixture so that a mass of the gas generant 8 used in the pellet 16 may remain as small as possible but yet still produce an effective amount of the inert gas mixture to extinguish the fire. A catalyst may also be present in the gas generant 8 to convert undesirable, toxic gases into less toxic, inert gases that may be used in fire suppression. The gaseous combustion products may be generated within a short amount of time after the gas generant 8 is ignited. For instance, the gas generant 8 may produce the gaseous combustion products within approximately 20 seconds to approximately 60 seconds after its ignition so that the inert gas mixture may be dispersed and the fire extinguished within approximately 30 seconds to approximately 60 seconds.

During combustion of the gas generant 8, substantially all of the combustion products that are solid at ambient temperature congeal into a solid mass, reducing particulates and smoke formed by combustion of the gas generant. The solid combustion products may produce a slag, which includes metallic elements, metal oxides, or combinations thereof. The slag may fuse on or near a burning surface of the pellet 16 when the gas generant 8 is combusted, producing a porous, monolithic frit. Since the slag fuses into a porous mass at or near the surface of the pellet 16 as it combusts, particulates produced during combustion of the pellet 16 may be minimized.

In one embodiment, the gas generant 8 is a HACN composition, as disclosed in United States Patent Nos. 5,439,537 and 6,039,820, both to Hinshaw *et al.* The HACN used in the gas generant 8 may be recrystallized and include less than approximately 0.1% activated charcoal or carbon. By maintaining a low amount of carbon in the gas generant 8, the amount of carbon-containing gases, such as CO, CO_2 , or mixtures thereof, may be minimized upon combustion of the gas generant 8. Alternatively, a

technical grade HACN having up to approximately 1% activated charcoal or carbon may be used. It is also contemplated that conventional gas generants 8 that produce gaseous combustion products that do not include carbon-containing gases or NO_x may also be used.

- 5 The HACN composition, or other gas generants 8, may include additional ingredients, such as at least one of an oxidizing agent, ignition enhancer, ballistic modifier, slag enhancing agent, cooling agent, chemical fire suppressant, inorganic binder, or an organic binder. Many additives used in the gas generant 8 may have multiple purposes. For sake of example only, an additive used as an oxidizer may
- 10 provide cooling, ballistic modifying, or slag enhancing properties to the gas generant 8. The oxidizing agent may be used to promote oxidation of the activated charcoal present in the HACN or of the ammonia groups coordinated to the cobalt in the HACN. The oxidizing agent may be an ammonium nitrate, an alkali metal nitrate, an alkaline earth nitrate, an ammonium perchlorate, an alkali metal perchlorate, an alkaline earth
- 15 perchlorate, an ammonium peroxide, an alkali metal peroxide, or an alkaline earth peroxide. The oxidizing agent may also be a transition metal-based oxidizer, such as a copper-based oxidizer, that includes, but is not limited to, basic copper nitrate ($[(\text{Cu}_2(\text{OH})_2\text{NO}_3)]$ ("BCN"), Cu_2O , or CuO . In addition to being oxidizers, the copper-based oxidizer may act as a coolant, a ballistic modifier, or a slag enhancing
- 20 agent. Upon combustion of the gas generant 8, the copper-based oxidizer may produce copper-containing combustion products, such as copper metal and cuprous oxide, which are miscible with cobalt combustion products, such as cobalt metal and cobaltous oxide. These combustion products produce a molten slag, which fuses at or near the burning surface of the pellet 16 and prevents particulates from being formed.
- 25 The copper-based oxidizer may also lower the pressure exponent of the gas generant 8, decreasing the pressure dependence of the burn rate. Typically, HACN-containing gas generants 8 that include copper-based oxidizers ignite more readily and burn more rapidly at or near atmospheric pressure. However, due to the lower pressure dependence, they burn less rapidly at extremely high pressures, such as those greater
- 30 than approximately 3000 psi (greater than approximately 20.68 MPa).

The ignition enhancer may be used to promote ignition of the gas generant 8 at a low positive pressure, such as from approximately 14 psi (approximately 0.097 MPa)

-10-

to approximately 500 psi (approximately 3.45 MPa). The ignition enhancer may be a conductive material having a large surface area. The ignition enhancer may include, but is not limited to, amorphous technical grade boron, high surface area flaked copper, or flaked bronze. The ballistic modifier may be used to decrease the burn rate pressure exponent of the gas generant. For instance, if the gas generant 8 includes cupric oxide and submicron particle size titanium dioxide, the gas generant may have a pressure exponent of less than approximately 0.3. Another ballistic modifier that may be used in the gas generant 8 is high surface area iron oxide. The ballistic modifier may also promote ignition of the gas generant 8. Additives that are able to provide ballistic modifying and ignition enhancing properties may include, but are not limited to, high surface area transition metal oxides and related species, such as basic copper nitrate and flaked metals, such as flaked copper.

The cooling agent may be used to lower the flame temperature of the gaseous combustion products. Since high flame temperatures contribute to the formation of toxic gases, such as NO and CO, cooling the gaseous combustion products is desirable. In addition, by using the cooling agent in the gas generant 8, less cooling of the gaseous combustion products may be necessary in the effluent train 6. The cooling agent may absorb heat due to its intrinsic heat capacity and, potentially, from an endothermic phase change, such as from a solid to a liquid, or an endothermic reaction, such as a decomposition of metal carbonates or metal hydroxides to metal oxides and carbon dioxide or water, respectively. Many of the additives previously described, such as the oxidizing agent, the ignition enhancer, and the ballistic modifier, may act as the cooling agent. For instance, the cooling agent may be a metal oxide, non-metal oxide, metal hydroxide, metal carbonate, or a hydrate thereof. However, desirably the cooling agent is not a strong oxidizing or reducing agent.

The slag enhancing agent may be used to meld the combustion products of the gas generant 8 into a cohesive solid, but porous, mass. Upon combustion of the gas generant 8, the slag enhancing agent may melt or produce molten combustion products that adhere to the solid combustion products and join the solid combustion products into the solid mass. Since the solid combustion products are melded together, the amount of smoke or particulates produced may be reduced. Silicon dioxide (SiO_2), titanium oxide, magnesium oxide, or copper-containing compounds may be used as the

-11-

slag enhancing agent. Desirably, titanium oxide or magnesium oxide is used because they produce low levels of NO_x upon combustion of the gas generant 8. The concentration of NO_x in the gaseous combustion products may also be reduced by including a catalyst for NO_x in the gas generant 8. For sake of example only, the catalyst may be tungsten oxide, which converts NO_x to nitrogen in the presence of ammonia.

The chemical fire suppressant or chemical fire retardant may also be used in the gas generant 8. The chemical fire suppressant may be a compound or a mixture of compounds that affects flames of the fire, such as a compound that delays ignition and reduces the spread of the flames in the space. The chemical fire suppressant may trap radicals, such as H, OH, O, or HO_2 radicals, which are important to oxidation in the vapor phase. The chemical fire suppressant may be a halogenated organic compound, a halogenated inorganic compound, or mixtures thereof.

The inorganic binder may provide enhanced pellet integrity when the pellet 16 is subjected to mechanical or thermal shock. The inorganic binder may be soluble in a solvent that is used to process the gas generant 8, such as water. As the solvent evaporates, the inorganic binder may coat solid particles of the gas generant 8, which enhances crush strength of granules and pellets 16 produced with the gas generant 8. In addition, since the binder is inorganic, carbon-containing gases such as CO or CO_2 , may not be produced when the gas generant is combusted. The inorganic binder may include, but is not limited to, a silicate, a borate, boric acid, or a mixture thereof. For instance, sodium silicate, sodium metasilicate ($\text{Na}_2\text{SiO}_3 \cdot 5\text{H}_2\text{O}$), sodium borosilicate, magnesium silicate, calcium silicate, aluminosilicate, aluminoborosilicate, or sodium borate may be used as the inorganic binder. In addition, HACN may act as the inorganic binder.

Small amounts of an organic binder may also be used in the gas generant 8 as long as minimal amounts of CO or CO_2 are produced during combustion. Gas generants 8 that include even a small amount of organic binder may have improved crush strength in pellet form compared to gas generants 8 that are free of organic binders. The organic binder may be present in the gas generant 8 from approximately 0.5% to approximately 2.0%. The organic binder may be a synthetic or naturally occurring polymer that dissolves or swells in water including, but not limited

-12-

- to, guar gum, polyacrylamide, and copolymers of polyacrylamide and sodium polyacrylate. The organic binder, in powder form, may be blended with dry ingredient(s) prior to the addition of water to promote dispersion of the organic binder. A sufficient amount of water may be added during mixing to produce a thick paste, which is subsequently dried and granulated prior to pelletization. Organic binders that dissolve or swell in organic solvents may also be used, such as ethyl cellulose, which dissolves or swells in ethanol. Gas generants 8 that include ethyl cellulose may be dry blended prior to mixing in the ethanol. The resulting thick paste may be subsequently dried and pressed into pellets 16. Curable polymeric resins may also be used as organic binders in the gas generant 8. The curable polymeric resin may be blended with the gas generant 8 and a curative in the absence of solvent or in the presence of a small amount of solvent to promote dispersion of the small amounts of the curable polymeric resin and the curative. The resulting powder may be pressed into a pellet 16 and allowed to cure at elevated temperature, such as at a temperature of approximately 135°F (approximately 57.2°C). The curable polymeric resin may include, but is not limited to, epoxy-cured polyesters and hydrosilylation-cured vinylsilicones. The organic binder may also include water-soluble, organic compounds that have a low carbon content, such as guanidine nitrate. If guanidine nitrate is used as the organic binder, it may be present in the gas generant 8 from approximately 1.0% to approximately 5.0%.

- In one embodiment, the gas generant 8 used in the fire suppression system 2 includes recrystallized HACN, cupric oxide (CuO), titanium dioxide (TiO_2), and high molecular weight polyacrylamide ($[\text{CH}_2\text{CH}(\text{CONH}_2)_n$). In another embodiment, the gas generant includes recrystallized HACN, CuO , silicon dioxide (SiO_2), TiO_2 , and polyacrylamide. In another embodiment, the gas generant includes recrystallized HACN, cuprous oxide (Cu_2O), and TiO_2 .

- The gas generant 8 may be produced by conventional methods, such as by using a vertical mixer, a muller mixer, a slurry reactor, or by dry blending the ingredients of the composition. In the vertical mixer, the solid ingredients of the gas generant 8 may be mixed in a solution that includes HACN dissolved in from approximately 15% by weight to approximately 45% by weight water. Ignitability and ease of combusting the gas generant 8 may increase when high concentrations of

-13-

HACN are dissolved during the mixing process. The water may be heated to 165°F (73.9°C) to increase the solubility of the HACN. Mixing the gas generant 8 at high water content (greater than approximately 35% by weight) and warm temperature (greater than approximately 145°F (greater than approximately 62.8°C)) dissolves at least a portion of the HACN and coats the additional ingredients. A high shear mixer, such as a dispersator, may be used to completely wet the high surface area solid ingredients before adding them to the vertical mixer or the high surface area solid ingredients may be preblended in a dry state. A powdered binder may be blended with the HACN prior to addition of water or another appropriate solvent. The slurry may be dried in a convection oven.

In one embodiment, a muller mixer is used to disperse the curable polymeric resin and the curative into the powdered ingredients of the gas generant 8. A small amount of solvent may also be added to promote dispersal of the curable polymeric resin and the curative. The gas generant 8 including the curable polymeric resin is allowed to cure once it has been pressed into the pellet 16.

To form the gas generant 8 in the slurry reactor, the HACN may be completely dissolved in water at a temperature of approximately 180°F (approximately 82.2°C). If technical grade HACN is used, any activated charcoal in the heated HACN solution may be removed, such as by filtration or another process. The heated HACN solution may be added to a cool, rapidly mixed suspension of the solid ingredients of the gas generant 8. Alternatively, a predispersed slurry of the solid ingredients may be slowly added to the rapidly stirred, HACN solution as it cools. Either of these methods may promote the formation of HACN crystallites on the insoluble solid ingredients of the gas generant 8. Once the suspension is cooled to a temperature ranging from at least approximately 80°F (approximately 26.7°C) to approximately 100°F (approximately 37.8°C), it may be filtered and the solids dried. The filtrate may be recycled as the liquid phase in subsequent slurry mixes.

To dry blend the gas generant 8, the HACN may be mixed with the other ingredients of the gas generant 8 using a v-shell, rotary cone, or Forberg blender. A small amount of moisture may be added to the mixture to minimize dusting. The mixture may then be dried before pelletization.

-14-

As previously described, the gas generant 8 or the igniter composition 14 and the gas generant 8 may be formed into the pellet 16. The pellet 16 may be formed by compressing the gas generant 8 or the igniter composition 14 and the gas generant 8 together to form a cylindrically-shaped pellet 16, as illustrated in FIG. 3a. However, the geometry of the gas generant 8 used in the fire suppression system 2 may depend on a desired ballistic performance of the gas generant 8, such as a desired burn rate or rate of evolution of the inert gas mixture as a function of time. Burn rates are typically categorized as a progressive burn, a regressive burn, or a neutral burn. A progressive burn is provided when the burning surface of the pellet 16 increases gradually as the pellet 16 burns. In a progressive burn, the rate of evolution of the inert gas mixture increases as a function of time. A regressive burn is provided when the burning surface of the pellet 16 decreases gradually as the pellet 16 burns. In a regressive burn, the rate of evolution of the inert gas mixture is initially high and decreases as a function of time. If the burning surface of the pellet 16 burns at a constant rate, a neutral burn is provided. In one embodiment, the gas generant 8 is formed into a pellet 16 having a center-perforated grain geometry, as illustrated in FIG. 3b. The center-perforated grain geometry has a high surface area, burns rapidly, and provides a neutral burn. The pellet 16 may also be formed into other shapes that provide a neutral burn as opposed to a regressive or progressive burn. The center-perforated pellet 16 may be produced using an appropriately designed die or by drilling a hole into a cylindrical pellet 16, using appropriate safety precautions.

The pellet 16 may include at least one layer of the igniter composition 14 in contact with one or more surfaces of the gas generant 8. A configuration of the igniter composition 14 used in the fire suppression system 2 may depend on the geometry of the gas generant 8. For instance, the pellet 16 may include a layer of the igniter composition 14 above a layer of the gas generant 8. Alternatively, a layer of the igniter composition 14 may be present below the gas generant 8 or may be present on multiple surfaces of the pellet 16. The igniter composition 14 may also be pressed on the surface of the pellet 16. Alternatively, the igniter composition 14 may be powdered, granulated, or pelletized and housed in a metal foil packet that is placed on or near the surface of the pellet 16. The metallic foil packet may include steel wool or another conductive material that absorbs heat from the igniter composition 14 and transfers it to

-15-

the surface of the gas generant 8. The igniter composition 14 may also be placed in a perforated flash tube within the center-perforation of the pellet 16. If the igniter composition 14 is granular or powdered, the perforated flash tube may be lined internally or externally with a metal foil or the igniter composition 14 may be inserted
5 into the perforated flash tube in preloaded foil packets.

In one embodiment, the igniter composition 14 includes from approximately 15% to approximately 30% boron and from approximately 70% to approximately 85% potassium nitrate. This igniter composition 14 is known in the art as "B/KNO₃" and may be formed by conventional techniques. In another embodiment,
10 an igniter composition 14 having strontium nitrate, magnesium, and small amounts of a polymeric organic binder, such as nylon, may be used. The igniter composition 14 is referred to herein as a Mg/Sr(NO₃)₂/binder composition. If the organic binder is nylon, the igniter composition 14 is referred to herein as a Mg/Sr(NO₃)₂/nylon composition. Since magnesium is water reactive, the organic binder used in the igniter
15 composition 14 may be soluble in organic solvents. For instance, ethyl cellulose or polyvinylacetate may also be used as the organic binder. The Mg/Sr(NO₃)₂/binder composition may be formed by conventional techniques. The igniter composition 14 may also include mixtures of B/KNO₃ and Mg/Sr(NO₃)₂/binder. The igniter compositions disclosed in United States Patent No. 6,086,693 may also be used as the
20 igniter composition 14.

The pellet 16 may be formed by layering the granules of the igniter composition 14 above or below the layer of the gas generant 8 in a die so that the igniter composition 14 and the gas generant 8 are in contact with one another. A pressure of approximately 8,000 psi (approximately 55.2 MPa) may be used to form
25 the pellet 16, which has a porosity ranging from approximately 5% to approximately 20%. The igniter composition 14 and the gas generant 8 may be compressed into the pellet 16 using a metal sleeve or a metal can, which provides support while the pellet 16 is being produced, handled, or stored. The metal can or the metal sleeve may also be used to inhibit burning of surfaces of the pellet 16 that are
30 enclosed by the metal sheathing. In the fire suppression system 2 of the present invention, the pellet 16 may burn at a controlled rate so that the amount of inert gas mixture produced during the burn remains constant as a function of time. To achieve a

-16-

neutral burn, at least one surface of the pellet 16 may be covered or inhibited by the metal can or metal sleeve so that these surfaces do not burn. An inner surface of the metal sheathing may also be painted with an inert inorganic material, such as sodium silicate or a suspension of magnesium oxide in sodium silicate, to inhibit the surfaces

5 of the pellet 16.

The pellets 16 may be housed in the combustion chamber 4 and have a total mass that is sufficient to produce an amount of the inert gas mixture sufficient for extinguishing the fire in the space. For sake of example only, in order to lower the oxygen concentration and extinguish a fire in a 1,000 cubic foot (28.32 cubic meter)

10 space, the gas generant 8 may have a total mass of approximately 40 pounds (approximately 18 kg). The inert gas mixture produced by the combustion of the gas generant 8 may lower the oxygen concentration in the space to a level that sustains human life for a limited duration of time. For instance, the oxygen concentration in the space may be lowered to approximately 13% by volume for approximately five

15 minutes

The combustion chamber 4 may be configured to house multiple pellets 16 of the gas generant 8 or the igniter composition 14 and the gas generant 8. Therefore, the fire suppression system 2 of the present invention may be easily configured for use in spaces of various sizes. For instance, the fire suppression system 2 may include one

20 pellet 16 if the fire suppression system 2 is to be used in a small space. However, if the fire suppression system 2 is to be used in a larger space, the combustion chamber 4 may include two or more pellets 16 so that the sufficient amount of the inert gas mixture may be produced. For sake of example only, in a 500 cubic foot (14.16 cubic meter) space, four pellets 16 having a 5.8-inch (14.73 cm) outer diameter, a 2.6-inch

25 (6.6 cm) height, and a weight of 4.44 pounds (2.01 kg) may be used, while eight of these pellets 16 may be used in a 1,000 cubic foot (28.32 cubic meter) space. In a 2,000 cubic foot (55.63 cubic meter) space, two generators, each containing eight pellets 16, may be strategically positioned. The pellets 16 may have an effective burning surface area so that the inert gas mixture may be produced within a short time

30 period after initiation of the gas generant 8. For instance, the inert gas mixture may be produced with approximately 20 seconds to approximately 60 seconds after initiation of the gas generant 8. If the fire suppression system 2 includes multiple pellets 16, the

-17-

pellets 16 may be ignited so that they are combusted simultaneously to provide a sufficient amount of the inert gas mixture to extinguish the fire. Alternatively, the pellets 16 may be ignited sequentially so that the inert gas mixture is produced at staggered intervals.

- 5 In one embodiment, the ignition train includes a squib, which, when electrically activated, ignites a granular or pelletized composition of B/KNO_3 in an ignition chamber. The hot effluents produced by combustion of the B/KNO_3 composition pass into the combustion chamber 4 and ignite the secondary ignition or igniter composition 14, which may be located in the metallic foil packet, pressed or painted on
10 the surface of the pellet 16, or placed in the perforated flash tube positioned in the center-perforation of the pellet 16.

- The fire suppression system 2 may be designed in various configurations depending on the size of the space in which the fire is to be extinguished. Exemplary configurations of the fire suppression system 2 include, but are not limited to, those
15 illustrated in FIGs. 1 and 4. As illustrated in FIG. 4, the fire suppression system 2 may have a tower configuration having a plurality of gas generators 70. A group or cluster of the gas generators 70 may be utilized to generate a sufficient amount of the inert gas mixture, which is delivered to the space in which the fire is to be suppressed. The number of gas generators 70 in the cluster and a controllable sequence in which the gas
20 generators 70 are initiated allows the ballistic performance of the fire suppression system 2 to be tailored to provide a sufficient amount of the inert gas mixture to the space. The number of gas generators 70 may also be adjusted to provide a desired mass flow rate history and action time of the inert gas mixture to the space. To configure the fire suppression system 2 for a particular space, gas generators 70 may be
25 added to or removed from the tower cluster. The fire sequencing used to initiate the gas generator 70 may be accomplished by controlling the timing of the electrical impulse to the initiating device 12 or by utilizing a pyrotechnic fuse. A column length of the pyrotechnic fuse may be selected to determine the time of initiation of the gas generator 70. The gas generator 70 may house the gas generant 8, which is illustrated
30 in FIG. 4 as having a center-perforated grain geometry. However, the gas generator 70 may accommodate other geometries of the gas generant 8 depending on the desired ballistic performance of the gas generant 8. The geometry of the igniter

-18-

composition 14 used in the fire suppression system 2 may depend on the grain geometry of the gas generant 8. For instance, the igniter composition 14 may be loaded into the metallic foil packets and placed on the surfaces of the gas generant 8. Alternatively, the igniter composition 14 may be placed in the perforated flash tube (not shown), which extends down the length of a center-perforated pellet 16 of the gas generant 8.

As previously described, the igniter composition 14 is ignited, which in turn combusts the gas generant 8 and produces the gaseous combustion products. The gaseous combustion products form the inert gas mixture, which then passes through a filter 18 and a controlling orifice 20 into a diffuser chamber 72. The filter 18 may be a screen mesh, a series of screen meshes, or a conventional filter device that removes particulates from the inert gas mixture. The filter 18 may also provide cooling of the inert gas mixture. The controlling orifice 20 may control the mass flow out of the gas generator 70 and, therefore, may control the flow rate of the inert gas mixture and the pressure within the gas generator 70. In other words, the controlling orifice 20 may be used to maintain a desired combustion pressure in the fire suppression system 2. The pressure in the gas generator 70 may be maintained at a level sufficient to promote ignition and to increase the burn rate of the gas generant 8. The pressure may also promote the reaction of reduced toxic gases, such as CO and NH₃, with gases that are oxidized, such as NO_x, which significantly reduces the concentration of these gases in the effluent gases. The controlling orifice 20 may be of a sufficient size to produce a combustion pressure ranging from approximately 600 psi (approximately 4.14 MPa) to approximately 800 psi (approximately 5.52 MPa) in the gas generator 70. Therefore, walls 22 of the gas generator 70 and of other portions of the fire suppression system 2 may be formed from a material that is capable of withstanding the maximum working pressure at the operating temperatures with appropriate engineering safety factors. In this tower configuration, high pressures are restricted to the small diameter, gas generator 70 volumes, while the remainder of the fire suppression system 2 operates at low pressures, which results in cost and weight savings.

In the diffuser chamber 72, plumes of the high velocity, inert gas mixture impinge on a flow deflector 74. The flow deflector 74 recirculates the inert gas mixture and results in a more uniform flow through a perforated diffuser plate or first

diffuser plate 24. The first diffuser plate 24 may disperse the inert gas mixture so that it does not exit the gas generator 70 as a high velocity jet. The inert gas mixture then passes through a heat management system 26 that includes cooling media or effluent scavenging media. The heat management system 26 may reduce the temperature of the inert gas mixture to a temperature that is appropriate to suppress the fire. Since combustion of the gas generant 8 produces a significant amount of heat in the gas generator 70, the inert gas mixture may be cooled before it is introduced into the space. For sake of example only, the heat released from a gas generant 8 combusted in a 2,000 cubic foot (56.63 cubic meter) space may be approximately 40,000 British Thermal Units ("BTU") (approximately 42,200,000 joules). In one embodiment, the heat management system 26 is a heat sink. The heat sink may be formed from conventional materials that are shaped into beds, beads, or tube clusters. The materials used in the heat sink may include, but are not limited to, metal, graphite, or ceramics. The material used in the heat sink and the geometry of the heat sink may be selected by one of ordinary skill in the art so that the heat sink provides the appropriate heat transfer surface, thermal conductivity, heat capacity, and thermal mass.

In another embodiment, the heat management system 26 includes a phase change material ("PCM"). The PCM removes thermal energy from the inert gas mixture by utilizing the PCM's latent heat of fusion and stores the thermal energy. The PCM may be an inert material that does not react with the inert gas mixture including, but not limited to, a carbonate, phosphate, or nitrate salt. For instance, the PCM may be lithium nitrate, sodium nitrate, potassium nitrate, or mixtures thereof. The PCM is described in more detail below.

The cooled, inert gas mixture may then be dispersed into the space through at least one final orifice 32, which reduces the pressure of the inert gas mixture relative to the pressure in the gas generator 70. The geometry of the final orifice(s) 32 may be selected based on the geometry of the space and the placement of the fire suppression system 2 in the space. Since the inert gas mixture is generated pyrotechnically, high pressure gas storage tanks and accompanying hardware to disperse the inert gas mixture may not be needed in the fire suppression system 2 of the present invention.

Another configuration of the fire suppression system 2 is shown in FIG. 1. The inert gas mixture, including nitrogen and water vapor, may be passed through the

-20-

filter 18 to remove any particulates that are produced upon combustion of the gas generant 8. The inert gas mixture may then be flowed through the controlling orifice 20 located at the exit of the combustion chamber 4. The controlling orifice 20 may control the mass flow out of the combustion chamber 4 and, therefore, may control the pressure within the combustion chamber 4. In other words, the controlling orifice 20 may be used to maintain a desired combustion pressure in the fire suppression system 2. The controlling orifice 20 may be of a sufficient size to produce a combustion pressure ranging from approximately 400 psi (approximately 2.76 MPa) to approximately 600 psi (approximately 4.14 Mpa) in the combustion chamber 4. Therefore, walls 22 of the combustion chamber 4 and of the effluent train 6 may be formed from a material capable of withstanding the maximum working pressure at the operating temperatures with appropriate engineering safety factors.

The combustion chamber 4 may also include the first diffuser plate 24 that disperses or diffuses the inert gas mixture into the heat management system 26 of the effluent train 6. The first diffuser plate 24 may disperse the inert gas mixture so that it does not exit the combustion chamber 4 as a high velocity jet. Rather, a laminar flow of the inert gas mixture may enter the effluent train 6. The effluent train 6 may include the heat management system 26 or a gas coolant material to reduce the temperature of the inert gas mixture to a temperature appropriate to suppress the fire. In one embodiment, the heat management system 26 is a heat sink, as previously described. In another embodiment, the heat management system 26 includes PCM 28. As previously described, the PCM 28 removes thermal energy from the inert gas mixture by utilizing the PCM's latent heat of fusion and stores the thermal energy. The PCM 28 may be an inert material that does not react with the inert gas mixture including, but not limited to, a carbonate, phosphate, or nitrate salt. For instance, the PCM 28 may be lithium nitrate, sodium nitrate, potassium nitrate, or mixtures thereof. The PCM 28 used in the heat management system 26 may be selected by one of ordinary skill in the art based on its phase change temperature, latent heat of fusion, or thermal properties, such as thermal conductivity, burn rate, heat capacity, density, or transition or melting temperature. In addition to these properties, the material selected as the PCM 28 may be dependent on the amount of time that is needed to ignite the gas generant 8 and produce the gaseous combustion products of the inert gas mixture. To

-21-

transfer heat from the inert gas mixture to the PCM 28, a tube cluster 30 may be embedded in, or surrounded by, the PCM 28. The tube cluster 30 may be formed from metal tubes that are capable of conducting heat, such as steel or copper tubes. The length, inner diameter, and outer diameter of the metal tubes may be selected by one of ordinary skill in the art depending on the amount of time required for the heat produced by the gas generant 8 to be conducted from the inert gas mixture to the PCM 28. The geometry of the tube cluster 30 in relation to the PCM 28 may be selected by one of ordinary skill in the art based on the amount of time necessary to ignite the gas generant 8 and produce gaseous combustion products and the amount of heat produced by the gas generant 8. When the inert gas mixture is flowed from the combustion chamber 4 and through the tube cluster 30, heat flux from the inert gas mixture may be transferred through the tube cluster 30 and into the PCM 28. When the PCM 28 is heated to its phase change temperature, it may begin to absorb its latent heat of fusion. Once the PCM 28 has absorbed its latent heat of fusion, an interface boundary temperature differential of the PCM 28 remains constant, which may enhance heat conduction from the surface of the tube cluster 30 to the PCM. Thermal energy may be stored in the PCM 28 based on the heat capacity of its liquid state once the PCM 28 has absorbed its latent heat of fusion.

The heat management system 26 may also be doped with a selective catalytic reduction ("SCR") catalyst or a non-selective catalytic reduction ("NSCR") catalyst to convert any undesirable gases that are produced as gaseous combustion products into gases that may be used in the inert gas mixture. For instance, the SCR and NSCR catalysts may be used to convert ammonia or nitrogen oxides into nitrogen and water, which may then be used in the inert gas mixture.

After the inert gas mixture has passed through the heat management system 26, the inert gas mixture may pass through a final orifice 32, which reduces the pressure of the inert gas mixture relative to the pressure in the combustion chamber 4. The inert gas mixture may then pass through a second diffuser plate 34 to uniformly disperse the inert gas mixture throughout the space. Since the inert gas mixture is generated pyrotechnically, high pressure gas storage tanks and accompanying hardware to disperse the inert gas mixture may not be needed in the fire suppression system 2 of the present invention.

-22-

The following are examples of gas generant compositions and igniter compositions for use within the scope of the present invention. These examples are merely illustrative and are not meant to limit the scope of the present invention in any way.

5

Examples

Example 1

A HACN Gas Generant Produced Using a Slurry Reactor

A gas generant including HACN, BCN, and Fe_2O_3 was produced in the slurry reactor. A 10 liter baffled slurry tank was filled with 4,900 grams of distilled water and stirred with a three blade stationary impeller at 600 revolutions per minute ("rpm"). A glycol heating bath was used to heat the water to 180°F (82.2°C). After the water temperature reached 180°F (82.2°C), 586.1 g of technical grade HACN was added to the mixer and stirred at 600 rpm for 10 minutes to allow the HACN to dissolve.

111.64 g of BCN and 18.56 g of Fe_2O_3 were dry blended together in a Nalgene™ quart container. 100 g of distilled water were then added into the blended BCN/ Fe_2O_3 and stirred for 5 minutes until an even suspension was made. 58 g of this suspension of BCN/ Fe_2O_3 /water was then injected slowly into the mix bowl with a 30 cc syringe while mixing rapidly. The slow addition of solid into the mix bowl allows for better oxidizer distribution in the mix. The heating system of the mix bowl was then turned off and the system was cooled at 1.4°F/minute (0.78°C/minute) by melting ice on the exterior of the mix bowl. When the mix temperature reached 160°F (71.1°C), a second addition of 58 g of BCN/ Fe_2O_3 /water was injected slowly into the mix bowl with a 30 cc syringe while mixing rapidly. Cooling with ice was continued after this addition.

When the temperature reached 139.7°F (59.83°C), a third addition of 58 g of BCN/ Fe_2O_3 /water was then injected slowly into the mix bowl with a 30 cc syringe while mixing rapidly. Cooling with ice was continued after this addition. When the temperature reached 119.9°F (48.83°C), 56.2 g (the remainder of the suspension) of BCN/ Fe_2O_3 /water was injected slowly into the mix bowl with a 30 cc syringe while mixing rapidly. Cooling with ice was continued after this addition until the temperature reached 75.4°F (24.1°C). At that time, the impeller was stopped and the material was transferred out of the mix bowl and into a five gallon bucket. The mix

-23-

was then filtered in a vacuum Erlenmeyer flask with a 1- μ m paper filter. The mixed gas generant was then placed onto a glass tray and dried at 165°F (73.9°C) overnight to remove any moisture.

5

Example 2

A HACN Gas Generant Produced By Vertical Mixing

A five gallon Baker Perkins vertical mixer was filled with 10,857 g of distilled water and stirred at 482 rpm. The mix bowl was heated to 165°F (73.9°C). After the water temperature reached 165°F (73.9°C), 3,160.0 g of recrystallized HACN was added into the mixer and stirred slowly at 482 rpm for 15 minutes to allow the HACN to partially dissolve and break up any clumps. 1,800 g of Cu_2O and 720 g of TiO_2 were then dry blended by sealing a five gallon bucket and shaking it. The mixer was stopped and the walls and blades were scraped down to incorporate any material that may have migrated up the mix blades. Then, the blend of Cu_2O and TiO_2 was added to the mix bowl and mixed for 15 minutes at 482 rpm. The mixer was stopped and the walls and blades were scraped down to incorporate any material that may have migrated up the mix blades. Then, 3,160 g of recrystallized HACN was added into the mix bowl and mixed for 15 minutes at 482 rpm. The mixer was stopped and the walls and blades were scraped down. The mixture was mixed for 30 minutes at 1,760 rpm. The mixer was stopped and the walls and blades were scraped. Then, the mixture was mixed for 30 minutes at 1,760 rpm. The mixture was loaded onto velo-stat lined trays and dried at 165°F (73.9°C). After drying, the coarse, granular material was granulated to a consistent small granule size using a Stokes granulator.

25

Example 3

A HACN Gas Generant with Organic Binder Produced By Vertical Mixing

To a one gallon Baker Perkins vertical mixer, 2,730 g of recrystallized HACN and 35 g of granular Cytec Cyanamer N-300 polyacrylamide were added. The two solids were blended for two minutes, after which 1,750 g of deionized water was added. The resulting slurry was mixed for 15 minutes. The mixer was stopped and the walls and blades were scraped down to incorporate any material that may have migrated up the mix blades.

30

-24-

In a two-gallon plastic container with a snap-on lid, 630 g of American Chemet Corp. UP13600FM cupric oxide and 105 g of DeGussa P-25 titanium dioxide were preblended by vigorous shaking. Then, the blend of cupric oxide and titanium dioxide was added into the mix bowl and mixed for 5 minutes. The mixer was stopped and the
5 walls and blades were scraped down to incorporate any material that may have migrated up the mix blades. The resulting paste was then mixed for an additional 15 minutes. The mixture was loaded into glass baking dishes and dried at 165°F (73.9°C) with occasional stirring. After drying, the coarse granular material was granulated to -12 mesh using a Stokes granulator.

10

Example 4

A HACN Gas Generant Produced in a Rotating Double-Cone Dryer

To a two cubic foot (0.057 cubic meter) rotating double-cone dryer, 2,996 g of cupric oxide and 817 g of titanium dioxide were added. The material was blended for
15 20 minutes by way of rotation of the rotating double-cone dryer. Afterwards, the inside walls of the rotating double-cone dryer were scraped down to free any unblended material. Next, 23,426 g of recrystallized HACN was added to the rotating double-cone dryer. The material was blended for an additional thirty minutes and then collected.

20

Example 5

A HACN Gas Generant Containing an Organic Binder Produced in a Muller Mixer

A polymer preblend was prepared by mixing 82 g of Crompton Corp. Fomrez F17-80 polyester resin with 17.4 g of Vantico Inc. Araldite MY0510 multifunctional
25 epoxy resin and 0.6 g of powdered magnesium carbonate. To a 12" (30.5 cm) diameter muller mixer, 10 g of the polymer preblend and 1,636 g of recrystallized HACN were added. This was blended for 10 minutes and the mixing surfaces were scraped down. Then, 294 g of American Chemet Corp. UP13600FM cupric oxide and 60 g of DeGussa P-25 titanium dioxide were added and the composition was mixed for
30 5 minutes. The mixer was again scraped down and the composition was blended for another 10 minutes. The composition was placed in a freezer and allowed to warm to room temperature immediately before pressing it into a pellet.

-25-

Example 6

Test Article Pellet Pressing

Pellets formed from the gas generants described in Examples 1, 2, or 4 were produced. To press the pellets, a 1.13 inch (2.87 cm) die assembly was used. A mold release agent, polytetrafluoroethylene ("PTFE"), was liberally applied to the die anvil and foot to minimize material sticking during the press cycle. 1.5 g of an igniter composition having a mixture of 60% B/KNO₃ and 40% Mg/Sr(NO₃)₂/binder was added to the die and leveled off with a spatula. The igniter composition was produced by blending together granules of the B/KNO₃ and Mg/Sr(NO₃)₂/binder. 10 g of the gas generant described in Examples 1, 2, or 4 was added to the die. The press foot was inserted into the top of the die assembly and twisted to ensure proper alignment. The pellet was pressed for 60 seconds at 8,000 lbf (35,590 N) and 8,000 psi (55.16 MPa). After pressing, the anvil was removed from the assembly and the pellet was pressed out of the die into a padded cup to minimize damage.

Example 7

Sleeved Test Article Pellet Pressing

Sleeved pellets formed from the gas generants described in Examples 1, 2, or 4 were produced. The press anvil and foot of the die were liberally sprayed with PTFE. A 1.05 inch (2.67 cm) internal diameter ("ID") steel ring was placed on the press anvil. 1.2 g of an igniter composition having a mixture of 60% B/KNO₃ and 40% Mg/Sr(NO₃)₂/binder was then added inside the steel ring. The surface of the igniter composition was then leveled with a spatula to ensure an even layer of the igniter composition on one surface of the pellet. An alignment sleeve was placed on top of the steel sleeve and 14.5 g of the gas generant described in Examples 1 or 2 was poured inside the alignment tool. A 1.00 inch (2.54 cm) outer diameter ("OD") press foot was inserted into the die. The sleeved pellet was pressed for 60 seconds at 6,900 lbf (30,690 N) and 8,000 psi (55.16 MPa). After pressing, the top surface of the sleeved pellet matched the top layer of the steel ring. Therefore, no post pressing process was required to remove the pellet from the press die. Instead, the anvil and alignment piece pulled off easily, leaving a filled steel ring of the gas generant.

-26-

Example 8

Sleeved Test Article Pellet Pressing with Hot Wire

Sleeved pellets were also pressed with embedded hot wires by running a loop of tungsten wire having a 0.010 inch (0.0254 cm) OD through two holes on the press anvil. The wire leads were rolled up and stored in the labeled opening on the underside of the press anvil. After installing the hot wire in the pressing fixture, the procedure for sleeved pellets (described in Example 7) was followed.

10

Example 9

5.8 Inch (14.7 cm) Diameter Test Pellets

3.3 pound (1.5 kg) pellets were pressed using a 150-ton (136,000 kg) hydraulic press. The anvil and press foot were sprayed liberally with PTFE. The anvil was then inserted into the die walls. 39.6 g of the igniter composition (40% B/KNO₃ and 60% Mg/Sr(NO₃)₂/binder) was added to the die by slowly pouring the material in a circular coil pattern starting at the center of the anvil and moving outward toward the die wall. The igniter composition was then leveled on top of the press anvil with a spatula. After ensuring an even layer of the igniter composition, 1,500 g of the gas generant described in Examples 1, 2, or 4 was added to the die. The press foot was then carefully inserted into the die. To ensure proper alignment, the press foot was spun around to ensure that no gas generant was trapped between the die walls and press foot. After alignment, the pellet was pressed at 211,000 lb_f (939,000 N) and 8,000 psi (55.16 MPa) for 60 seconds. To remove the pellet, the press anvil was removed and the die walls were positioned on top of a 6.0 inch (15.2 cm) inner diameter ("ID") knockout cup. A slight amount of force was applied to the press foot to push the pellet out of the 5.8 inch (14.7 cm) die walls.

30

Example 10

Test Pellets Pressed in a Steel Can

The gas generant (737 g) described in Example 4 was added to a carbon steel can having an OD of 6.0 inches (15.2 cm), an ID of 5.8 inches (14.7 cm), a height of 2.15 inches (5.46 cm), and a depth of 2.06 inches (5.23 cm) and pressed using a

-27-

150-ton (136,000 kg) hydraulic press to a maximum pressure of 8,042 psi (55.45 MPa). Pressure was maintained at or above 8,000 psi (55.16 MPa) for one minute. A second addition of 740 g of the gas generant was added to the press die along with a 59.4 g blend of an igniter composition that included 11% B/KNO₃ and 89%

- 5 Mg/Sr(NO₃)₂/binder. The igniter composition was spread evenly on the top surface of the gas generant. The remaining gas generant and the igniter composition were then pressed at 8,197 psi (56.52 MPa) for one minute. The total height of the gas generant and igniter composition after the final press cycle was 2.01 inches (5.11 cm).

10

Example 11

Subscale Fire Suppression System

- A subscale system of the fire suppression system 2 was produced, as shown in FIG. 2. The gas generant 8 used in the subscale system included a composition of HACN, Cu₂O, and TiO₂, which was prepared as previously described. The igniter
15 composition 14 included 1 g of 60% B/KNO₃ and 40% Mg/Sr(NO₃)₂/binder. The subscale system included an igniter cover 36, an inner case 40, an outer case 42, a base 44, a perforated tube 46, a screen retainer 48, a cover fabrication 50, an inner barrier 52, a tie rod 54, a perforated baffle 56, a boss 58, and a baffle 60. An inhibitor 62, formed from Krylon/Tape, was applied to the bottom of the gas generant
20 pellet 16, which came in contact with a spacer 64 in the combustion chamber 4. In addition to providing heat management properties, the perforated tube 46 prevents the escape of particulates from the ignition chamber.

- The mass of the gas generant 8 in the fire suppression system 2 was selected so that when the inert gas mixture was vented into a 100 cubic foot (2.83 cubic meter)
25 enclosure, atmospheric oxygen was displaced and removed to a level low enough to extinguish combustion in the enclosure. A 3.3 lb (1.5 kg) pellet having the gas generant 8 was used in the subscale system. Upon combustion of the pellet, the oxygen content in the 100 cubic foot (2.83 cubic meter) enclosure was reduced to below approximately 13% oxygen, as shown in FIG. 5.

- 30 In test A, a cylindrical pellet 16 was tested. The pressure generated in the combustion chamber 4 and the temperature of the gas in the aft of the combustion chamber 4 were measured. As shown in FIG. 6, the maximum pressure in the fire

-28-

suppression system 2 was slightly more than 300 psi (2.07 MPa) at approximately 9 seconds after ignition of the gas generant 8. The maximum temperature in the fire suppression system 2 was less than 500°F (260°C) at approximately 9 seconds after ignition of the gas generant 8.

- 5 In test B, a cylindrical pellet that was pressed into a metal cylinder and inhibited on one end was tested. As shown in FIG. 7, the maximum pressure in the fire suppression system 2 was approximately 650 psi (approximately 4.48 MPa) at approximately 18 seconds after ignition of the gas generant 8. The maximum temperature in the fire suppression system 2 was less than approximately 550°F (less
10 than approximately 288°C) at approximately 19 seconds after ignition of the gas generant 8.

Example 12

Mini-Generator Test

- 15 A mini-generator developed for use in airbag research was used to test pellets of the igniter composition 14 and gas generant 8 described in Examples 6 or 7. The mini-generator is a conventional device that consists of reuseable hardware and is a simplified prototype of a driver-side airbag inflator.

- Pellets 16 having a mass of from approximately 20 g to approximately 25 g
20 were ignited in the mini-generator. The gaseous combustion products (or effluent gases) of the pellets 16 were transferred into gas-impermeable bags and tested to determine the contents of the gaseous combustion products. The gaseous combustion products were tested using a conventional, colorimetric assay, i.e., the Draeger Tube System, which is known in the art. In the mini-generator, CO levels decreased from
25 2,000 parts per million ("ppm") to 50 ppm. NO_x levels decreased from 2,000 ppm to 150 ppm. In addition, a tough, unitary slag was produced.

Example 13

100 Cubic Foot (2.83 Cubic Meter) Tank Test

- 30 The pellets 16 described in Example 10 were tested in the subscale fire suppression system described in Example 11, which was attached vertically to an assembly plate near the bottom of a 100 cubic foot (2.83 cubic meter) test tank

-29-

equipped with pressure transducers, thermocouples, a video camera, and an oxygen sensor. The tank was designed with a vent to eliminate significant overpressure. A Thiokol ES013 squib was electronically activated and the hot effluents produced by the squib ignited 6 grams of B/KNO₃ in the ignition chamber, which in turn ignited the igniter composition 14 that was pressed onto the top surface of the gas generant 8. The igniter composition 14 then ignited the gas generant 8. The pressure in the combustion chamber reached a maximum pressure of 650 psi (4.48 MPa) in about 18 seconds. The pressure in the combustion chamber decreased to 50 psi (0.35 MPa) 25 seconds after ignition. Maximum pressure in the 100 cubic foot (2.83 cubic meter) tank was 0.024 psig (166 Pa). After the test, ammonia, carbon monoxide, NO_x, and nitrogen dioxide were measured using appropriate Draeger tubes at 48 ppm, 170 ppm, 105 ppm and 9 ppm, respectively.

Example 14

Use of Igniter Composition Placed on the Surface of the Gas Generant Grain

A pellet 16 was pressed into a can similarly to that described in Example 10, except that the igniter composition was not pressed onto the top surface of the gas generant 8. When the resulting pellet 16 was tested in the subscale fire suppression system described in Example 11, the Thiokol ES013 squib ignited 1 g of B/KNO₃ in the ignition chamber which, in turn, ignited a 59.4 g blend of the igniter composition (11% B/KNO₃ and 89% Mg/Sr(NO₃)₂/binder) assembled in an aluminum foil packet placed on the top surface of the gas generant 8. Ignition was enhanced over that obtained in Example 13 because the maximum pressure of 900 psi (6.21 MPa) in the combustion chamber was reached at 16 seconds after ignition.

Example 15

Use of Flaked Copper-Containing Metals as an Ignition Aid

Two 10 g, 1.1-OD cylindrical pellets 16 were pressed at 8,000 psi (55.16 MPa). One pellet 16 included the gas generant 8 described in Example 4. The other pellet 16 included 90% by weight of the gas generant 8 described in Example 4 blended with 10% by weight of Warner-Bronz finely-divided bronze flakes, produced by Warner Electric Co., Inc. On the top surface of each pellet 16, 0.5 g of granular

-30-

Mg/Sr(NO₃)₂/binder was present. The igniter composition 14 on each pellet 16 was ignited by a hot wire. The pellet 16 that included the finely divided bronze flakes ignited more smoothly, combusted more rapidly, and produced a stiffer slag once combusted compared to the pellet 16 without the finely divided bronze flakes.

5

Example 16

Evaluation of Binders in HACN Gas Generants (Small Scale)

HACN gas generant compositions were mixed similarly to those described in Examples 2, 3, 4, and 5. For each composition, three 0.5 inch (1.27 cm) diameter, 4.0 g pellets were pressed at 2,000 lbs force (8,900 N) for 20 seconds. In addition, three 1.1 inch (2.79 cm) diameter, 15.0 g pellets were pressed at 10,000 lbs force (44,500 N) for 20 seconds. The pellets were analyzed for crush strength at a 0.125 in/min (0.318 cm/min) compression rate. The 0.5 inch (1.27 cm) pellets were used to determine axial crush strength and the 1.1 inch (2.79 cm) diameter pellets were analyzed for radial crush strength. The data are summarized in Table 1 and show that pellets 16 having the organic binder or inorganic binder had improved axial crush strength compared to those compositions having no binder. In addition, many of the pellets 16 had improved radial crush strength compared to those compositions having no binder.

20

Table 1: Crush Strength of HACN Gas Generants^a as a Function of Binder.

Binder	% HACN	% CuO	Mix Method (Ex. #)	Pellet Density (g/cc)	Axial Crush Strength (lbs/kg)	Radial Crush Strength (lbs/kg)
None	86.0	11.0	4	1.751	319/145	65/30
None	86.0	11.0	2	1.753	296/134	123/55.8
0.5% cured polyester	81.8	14.7	5	1.841	417/189	121/54.9
1.0% cured polyester	77.7	18.3	5	1.900	610/277	182/82.6
2.0% cured polyester	69.3	25.7	5	2.020	795/361	253/115
3.0% cured polyester	61.0	33.0	5	2.17	1059/480	365/166

-31-

Binder	% HACN	% CuO	Mix Method (Ex. #)	Pellet Density (g/cc)	Axial Crush Strength (lbs/kg)	Radial Crush Strength (lbs/kg)
2.0% guar	74.5	20.5	5	1.812	757/343	178/80.7
1.0% polyacrylamide	78.0	18.0	4	1.751	507/230	220/99.8
1.5% polyacrylamide	74.1	21.4	4	1.789	574/260	210/95.3
2.0% polyacrylamide	70.1	24.3	4	1.819	586/266	245.7/111.5
1.5% copolymer ^b	78.0	17.5	4	1.792	672/305	232/105
4.0% guanidine nitrate	79.2	13.8	4	1.762	373/169	149/67.6
1.0% ethyl cellulose	77.0	19.0	4	1.836	609/276	181/82.1
1.5% cured silicone	71.4	23.8	5	1.949	336/152	46/20.9
2.5% sodium silicate	84.1	10.4	4	1.725	403/183	217/98.4

^a All formulations include 3% titanium dioxide.

^b The copolymer includes 90% sodium acrylate and 10% acrylamide monomers, respectively.

- Gas-generator hardware larger in scale than that used in Example 17 was used
- 5 to test 1.42 inch (3.61 cm) diameter pellets 16 of formulations selected from Table 1. The 1.42 inch (3.61 cm) diameter pellets were produced by pressing 58.0 g of the gas generant at 16,000 lbs force (71, 200 N) for 60 seconds. Behind a protective shield, a hole was drilled into the center of each of the pellets 16 using a 0.3015 inch (0.7658 cm) OD drill bit to produce a center-perforation in the pellets. The gas generator
- 10 hardware was attached to a 60-liter tank. The pellets were then ignited and combustion analyses were performed on the gaseous combustion products. After combustion, dilution of the air in the 60-liter tank by combustion gases produced by the gas generant 8 was sufficient to decrease oxygen content in the tank to approximately 13%. Results of these combustion analyses are summarized in Table 2.

Table 2: Combustion Analysis of Small Center-Perforated Gas Generant Pellets

Binder	Test Info ¹	Pellet Density (g/cc)	Maximum Pressure (psi/MPa)	Rise Time (sec)	NH ₃ (ppm)	NO _x (ppm)	CO (ppm)	NO ₂ (ppm)
dry blended, no binder	1a	1.664	690.4/4.760	1.10	7	55	230	17
dry blended, no binder	2a	1.728	688.5/4.747	2.16	86	85		12
wet mixed, no binder	1a	1.668	584.0/4.027	1.28	85	80	220	17
1% polyacrylamide	1a	1.764	402.3/2.774	2.21	5	105	850	60
1% polyacrylamide	2a	1.762	528.3/3.643	1.00	83	90	850	28
2% guar	1b	1.674	637.7/4.397	0.92	170	55	1900	2
1% cured polyester	1a	1.875	800.8/5.521	1.50	40	85	680	60
1% ethyl cellulose	1a	1.829	390.6/2.693	1.97	10	150	1200	85
1% copolymer ²	1a	1.769	254.9/1.758	3.76	23	300	1200	150
4% guanidine nitrate	2a	1.737	752.9/5.191	1.05	58	70	1700	12
2.5% sodium silicate	2b	1.706	1299.8/8.962	11.63	340	125	380	40
1.5% silicone	2a	1.945	1391.6/9.595	10.14			1100	

¹ (1) Signifies the use of 1 g of B/KNO₃ in the ignition chamber, 1 g of Mg/Sr(NO₃)₂/binder in an aluminum foil packet on top of the pellet, and 1 g of Mg/Sr(NO₃)₂/binder in the pellet's center perforation; (2) Signifies the use of 1 g of B/KNO₃ in the ignition chamber and 2 g of Mg/Sr(NO₃)₂/binder in an aluminum foil packet on top of the pellet; (a) Signifies the combustion chamber limiting orifice diameter of 0.086" (0.218 cm); (b) Signifies an orifice diameter of 0.0785" (0.199 cm).

² The copolymer includes 90% sodium acrylate and 10% acrylamide monomers, respectively.

Example 17

Evaluation of Binders in HACN Gas Generants (Larger Scale)

Larger, center-perforated pellets were fabricated by pressing 1,520 g of the HACN gas generant 8 in a 5.8" (14.7 cm) diameter die at 8,000 psi (55.16 MPa) for a

-33-

- minimum of 1 minute. Once the pellets were pressed, a 1.25" (3.18 cm) diameter drill bit was used to produce a center perforation in the pellets. The pellets were tested in fire suppression system 2 as illustrated in FIG. 2 using the 100 cubic foot (2.83 cubic meter) tank test described in Example 13. The ignition train utilized an ATK Thiokol
- 5 Propulsion ES013 squib, 2 g of B/KNO₃ in the ignition chamber and 50 g of Mg/Sr(NO₃)₂/binder igniter composition in a foil packet placed on top of the center-perforated pellet. The pellets were then ignited and combustion analyses were performed on the gaseous combustion products. The combustion analyses are summarized in Table 3. Measured toxic gaseous effluent levels were generally lower
- 10 in the larger scale tests compared to those in the small scale tests, which were described in Example 16.

Table 3: Larger Scale Gas Generant Combustion Analysis Tests¹

Binder	Limiting Orifice Diameter (in)	Pellet Density (g/cc)	Maximum Pressure (psi/MPa)	Rise Time (sec)	NH ₃ (ppm)	NO _x (ppm)	CO ₂ (%)	CO (ppm)
dry blended, no binder	9/32	1.792 ²	913.0/6.295	2.70	35	33		29
wet mixed, no binder	9/32	-	787.0/5.426	2.50	40	40		23
0.5% cured polyester	9/32	1.827	684.0/4.716	3.37	48	45	0.22	175
4.0% guanidine nitrate	5/16	1.732	657.7/4.535		18	42	0.32	300
4.0% guanidine nitrate	5/16	1.719	553.7/3.818	3.33	35	47	0.28	300
1.0% polyacrylamide	9/32	1.724	543.0/3.744	2.68	8	25	0.30	270
1.0% polyacrylamide	9/32	1.727	542.0/3.737	2.50	7	23		265
1.0% polyacrylamide using HACN co-crystallized with 0.9% charcoal (tech. grade)	9/32	1.750	484.9/3.343	2.61	9	45		840

-34-

Binder	Limiting Orifice Diameter (in)	Pellet Density (g/cc)	Maximum Pressure (psi/MPa)	Rise Time (sec)	NH ₃ (ppm)	NO _x (ppm)	CO ₂ (%)	CO (ppm)
HACN) ³								
0.5% polyacrylamide using tech. grade HACN ⁴	9/32	1.735	572.0/3.944	2.50	11	60	0.62	670
1.0% polyacrylamide using 50% tech. grade HACN ⁵	9/32	1.865	412.0/2.841	3.60	11	45	0.61	670

¹ Nitrogen dioxide was not detected in these tests using Draeger tubes and, thus, nitrogen dioxide is assumed to be less than 1 ppm. Unless noted otherwise, recrystallized HACN was used in the compositions tested.

² Pellet pressed at 11,000 psi (75.84 MPa).

5 ³ Formulation includes 71% tech. grade HACN, 25% cupric oxide and 3% titanium dioxide.

⁴ Formulation includes 74.5% tech. grade HACN, 22% cupric oxide and 3% titanium dioxide.

⁵ Formulation includes 37.2% carbon-free HACN, 37.2% tech. grade HACN, 21.6% cupric oxide and 3% titanium dioxide.

10 While the invention may be susceptible to various modifications and alternative forms, specific embodiments have been shown by way of example in the drawings and have been described in detail herein. However, it should be understood that the invention is not intended to be limited to the particular forms disclosed. Rather, the invention is to cover all modifications, equivalents, and alternatives falling within the

15 spirit and scope of the invention as defined by the following appended claims.

-35-

CLAIMS

What is claimed is:

- 5 1. A fire suppression system, comprising:
a gas generant formulated to pyrotechnically produce an inert gas mixture suitable for
extinguishing a fire; and
a heat management system.
- 10 2. The fire suppression system of claim 1, wherein the gas generant is
present in a combustion chamber of the fire suppression system.
3. The fire suppression system of claim 1, wherein the gas generant is
present in the fire suppression system in at least one pellet.
- 15 4. The fire suppression system of claim 3, wherein the at least one pellet is
formed into a geometry that provides a neutral burn when combusted.
5. The fire suppression system of claim 3, wherein the at least one pellet
20 has a total mass sufficient to produce an amount of the inert gas mixture sufficient to
extinguish the fire.
6. The fire suppression system of claim 1, further comprising an igniter
composition in contact with the gas generant.
- 25 7. The fire suppression system of claim 6, wherein the igniter composition
is formulated and of sufficient mass to produce an amount of heat sufficient to ignite
the gas generant.
- 30 8. The fire suppression system of claim 6, wherein the igniter composition
is formulated to produce solid combustion products when combusted.

-36-

9. The fire suppression system of claim 6, wherein the igniter composition and the gas generant are compressed together in the at least one pellet.

10. The fire suppression system of claim 6, wherein the igniter composition comprises from approximately 15% to approximately 30% boron and from approximately 70% to approximately 85% potassium nitrate.

11. The fire suppression system of claim 6, wherein the igniter composition comprises strontium nitrate, magnesium, and an organic binder.

12. The fire suppression system of claim 1, wherein the gas generant is formulated to produce at least one gaseous combustion product and at least one solid combustion product when combusted.

13. The fire suppression system of claim 12, wherein the inert gas mixture is formed from substantially all of the at least one gaseous combustion product produced by combustion of the gas generant.

14. The fire suppression system of claim 12, wherein the at least one solid combustion product is formulated to minimize production of particulates during combustion of the gas generant.

15. The fire suppression system of claim 12, wherein the at least one solid combustion product produced by combustion of the gas generant is a slag.

16. The fire suppression system of claim 15, wherein the slag is present on a surface of the at least one pellet.

17. The fire suppression system of claim 1, wherein the gas generant is formulated to produce minimal amounts of carbon monoxide, particulates, or smoke when combusted.

-37-

18. The fire suppression system of claim 1, wherein the gas generant is formulated to produce less than an Immediately Harmful to Life or Health value of ammonia, carbon monoxide, carbon dioxide, or nitrogen oxides when combusted.
- 5 19. The fire suppression system of claim 1, wherein the gas generant is formulated to produce less than 1 percent of an original weight of the gas generant in particulates or smoke.
20. The fire suppression system of claim 1, wherein the inert gas mixture
10 comprises nitrogen and water.
21. The fire suppression system of claim 1, wherein the gas generant comprises an oxidizer, a fuel, and a binder.
- 15 22. The fire suppression system of claim 1, wherein the gas generant further comprises at least one of an oxidizing agent, an ignition enhancer, a ballistic modifier, a slag enhancing agent, a cooling agent, and a binder.
23. The fire suppression system of claim 1, wherein the gas generant
20 comprises hexa(amine)cobalt(III)-nitrate, cuprous oxide, and titanium dioxide.
24. The fire suppression system of claim 1, wherein the gas generant comprises hexa(amine)cobalt(III)-nitrate, cupric oxide, titanium dioxide, and polyacrylamide.
- 25 25. The fire suppression system of claim 1, wherein the heat management system is configured to reduce the temperature of the inert gas mixture.
26. The fire suppression system of claim 1, wherein the heat management
30 system comprises a heat sink or a phase change material.

-38-

27. The fire suppression system of claim 26, wherein the phase change material comprises lithium nitrate, sodium nitrate, potassium nitrate, or mixtures thereof.
- 5 28. The fire suppression system of claim 26, wherein the fire suppression system is configured to transfer heat from the inert gas mixture to the phase change material.
- 10 29. The fire suppression system of claim 1, wherein the fire suppression system is configured to disperse the inert gas mixture therefrom within from approximately 20 seconds to approximately 60 seconds after ignition of the gas generant.
- 15 30. The fire suppression system of claim 1, further comprising at least one diffuser plate to disperse the inert gas mixture.
31. The fire suppression system of claim 30, wherein the at least one diffuser plate is configured and positioned to diffuse the inert gas mixture into the heat management system or is configured and positioned to disperse the inert gas mixture exiting from the fire suppression system.
- 20 32. A method for fighting a fire in a space, comprising:
igniting a gas generant to produce an inert gas mixture that comprises minimal amounts of carbon monoxide, particulates, or smoke; and
25 introducing the inert gas mixture into a space.
33. The method of claim 32, wherein igniting a gas generant to produce an inert gas mixture comprising minimal amounts of carbon monoxide, particulates, or smoke comprises igniting the gas generant to produce nitrogen and water.

30

-39-

34. The method of claim 32, wherein igniting a gas generant to produce an inert gas mixture comprises igniting a nonazide gas generant composition to produce gaseous combustion products and solid combustion products.
- 5 35. The method of claim 34, wherein igniting a gas generant to produce an inert gas mixture comprises forming the inert gas mixture with substantially all of the gaseous combustion products produced by igniting the gas generant.
36. The method of claim 34, wherein igniting a gas generant to produce an
10 inert gas mixture comprises generating the gaseous combustion products within from approximately 20 seconds to approximately 60 seconds after ignition of the gas generant.
37. The method of claim 34, wherein igniting a gas generant to produce an
15 inert gas mixture comprises producing the gaseous combustion products that are substantially free of carbon-containing gases or nitrogen oxides.
38. The method of claim 32, wherein igniting a gas generant to produce an
20 inert gas mixture comprises producing a neutral burn of the gas generant.
39. The method of claim 32, wherein igniting a gas generant to produce an inert gas mixture comprises igniting an igniter composition to produce sufficient heat to ignite the gas generant.
- 25 40. The method of claim 39, wherein igniting an igniter composition to produce sufficient heat to ignite the gas generant comprises igniting an igniter composition comprising from approximately 15% to approximately 30% boron and from approximately 70% to approximately 85% potassium nitrate.
- 30 41. The method of claim 39, wherein igniting an igniter composition to produce sufficient heat to ignite the gas generant comprises igniting an igniter composition comprising strontium nitrate, magnesium, and an organic binder.

42. The method of claim 32, wherein igniting a gas generant to produce an inert gas mixture comprises producing solid combustion products that minimize the particulates and the smoke formed by igniting the gas generant.

5

43. The method of claim 32, wherein igniting a gas generant to produce an inert gas mixture comprises igniting a gas generant that comprises hexa(ammine)cobalt(III)-nitrate, cuprous oxide, and titanium dioxide.

10

44. The method of claim 32, wherein igniting a gas generant to produce an inert gas mixture comprises igniting a gas generant that comprises hexa(ammine)cobalt(III)-nitrate, cupric oxide, titanium dioxide, and polyacrylamide.

15

45. The method of claim 32, wherein introducing the inert gas mixture into a space comprises dispersing the inert gas mixture into the space within from approximately 20 seconds to approximately 60 seconds after ignition of the gas generant.

20

46. The method of claim 32, further comprising reducing a temperature of the inert gas mixture after combustion of the gas generant.

25

47. The method of claim 46, wherein reducing a temperature of the inert gas mixture after combustion of the gas generant comprises exposing the inert gas mixture to a heat management system.

25

48. The method of claim 47, wherein exposing the inert gas mixture to a heat management system comprises flowing the inert gas mixture into a heat sink or flowing the inert gas mixture over a phase change material.

30

49. The method of claim 32, further comprising extinguishing the fire by reducing an oxygen content in the space.

-41-

50. The method of claim 49, wherein extinguishing the fire by reducing an oxygen content in the space comprises reducing the oxygen content to approximately 13% by volume.

1/5

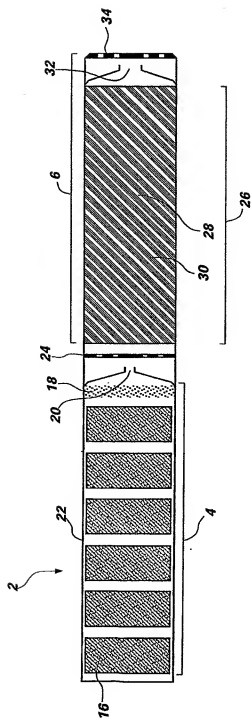


FIG. 1

2/5

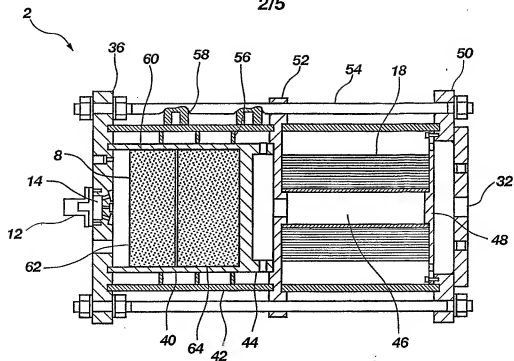


FIG. 2

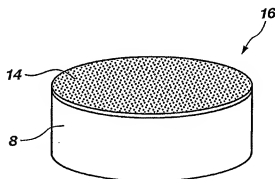


FIG. 3a

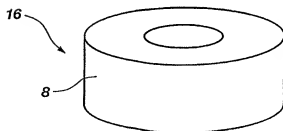


FIG. 3b

3/5

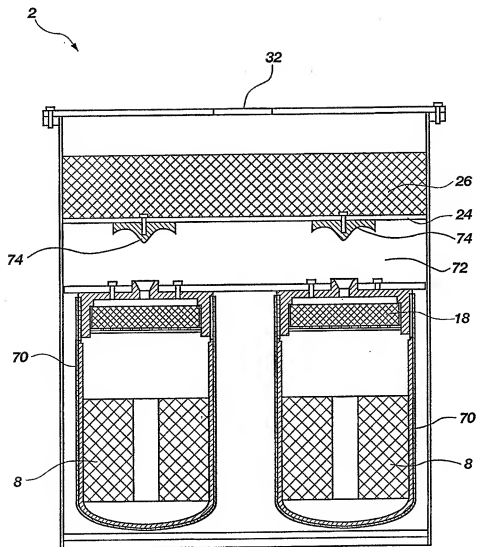


FIG. 4

4/5

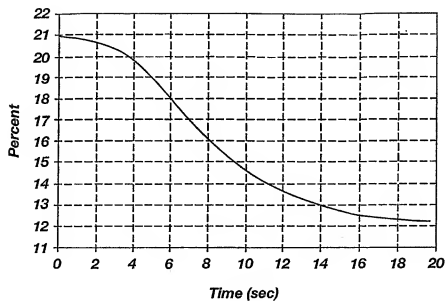


FIG. 5

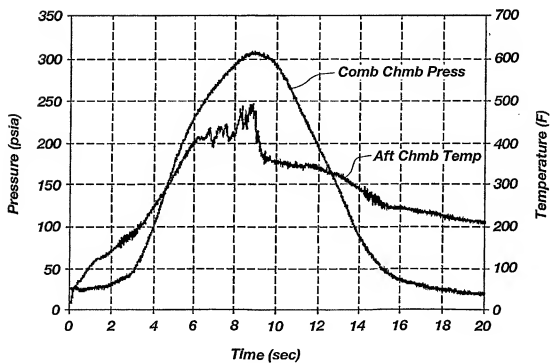
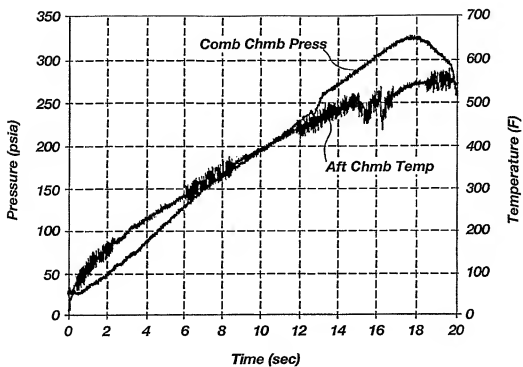


FIG. 6

5/5

**FIG. 7**